# Evolution of helimagnetic correlations when approaching the quantum critical point of $Mn_{1-x}Fe_xSi$

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We present a comprehensive investigation of the evolution of helimagnetic correlations in  $Mn_{1-x}Fe_xSi$  with increasing doping. By combining polarized neutron scattering and high resolution neutron spin echo spectroscopy we investigate three samples with x = 0.09, 0.11, and 0.14, i.e., with compositions on both sides of the concentration  $x^* \sim 0.11$  where the helimagnetic Bragg peaks disappear and between  $x^*$  and the quantum critical concentration  $x_c \sim 0.17$ , where  $T_c$  vanishes. We find that the abrupt disappearance of the long range helical periodicity at  $x^*$  does not affect the precursor fluctuating correlations. These build up with decreasing temperature in a similar way as for the parent compound MnSi. Also the dynamics bears strong similarities to MnSi. The analysis of our results indicates that frustration, possibly due to achiral Ruderman-Kittel-Kasuya-Yosida (RKKY) interactions, increases with increasing Fe doping. We argue that this effect explains both the expansion of the precursor phase with increasing x and the abrupt disappearance of long range helimagnetic periodicity at  $x^*$ .

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### I. INTRODUCTION

The physics of the chiral magnet MnSi touches several fundamental questions in condensed-matter physics, from the stabilization of exotic states like chiral skyrmions [1] to the interplay between localized and itinerant magnetism [2-8] as well as to non-Fermi-liquid behavior [9–14] and quantum fluctuations [15–23]. Under pressure a non-Fermi-liquid behavior sets in without quantum criticality [11,15,24,25]. Furthermore, the first-order transition temperature  $T_C$  is driven to 0 K at  $p_C \sim 1.4$  GPa, although the magnetic moment does not vanish. In the region of the temperature-pressure phase diagram where  $p \gtrsim p_C$  and  $T_C = 0$  K, long range range spiral and skyrmion correlations are restored under magnetic fields [7], a result that has been attributed to a softening of the magnetic moment. Pressure would therefore enhance the itinerant electron character of magnetism, triggering the suppression of  $T_C$ . On the other hand, in the absence of quantum critical point (QCP) at  $p_C$  [19], the origin and nature of the highly debated non-Fermi-liquid phase [16–18,20,26,27] remains an open question. It was suggested that this phase, out of which magnetic fields induce long range spiral correlations, is fluctuating and possibly of quantum nature [19]. However, our high resolution neutron spin echo (NSE) spectroscopy measurements did not reveal the existence of such fluctuations, possibly due to limitations (background contribution of the pressure cell) inherent to measurements under high pressures [7].

These limitations are overcome by chemical pressure, in the form of Fe doping in  $Mn_{1-r}Fe_rSi$ . The behavior of this system resembles that of MnSi under pressure [28-37], and our magnetization, susceptibility, and SANS investigations [32,33] led to the phase diagram shown in Fig. 1. With increasing doping, the transition temperature decreases continuously and vanishes at  $x_C \sim 0.17$ . On the other hand, the helimagnetic Bragg peaks, a signature of long range helimagnetic periodicity, disappear abruptly at a much lower concentration of  $x^* \sim 0.11$ . For  $x \gtrsim x^*$  magnetic susceptibility and electric transport phenomena reveal a non-Fermi-liquid behavior, which, as in MnSi under pressure, has been attributed to a chiral spin liquid state governed by quantum fluctuations [35-37]. Here we investigate the evolution of helimagnetic correlations and their dynamics as a function of chemical substitution in  $Mn_{1-x}Fe_xSi$  using polarized neutron scattering and neutron spin echo (NSE) spectroscopy. We investigated three samples with x = 0.09, 0.11 and 0.14, i.e., with compositions on both sides of  $x^*$  and between  $x^*$  and  $x_{\rm C}$ . Our results reveal that the abrupt disappearance of long range helimagnetic periodicity at  $x^*$  does not affect the precursor fluctuating correlations. These build up with decreasing temperature in a way that is very similar to that of the parent compound MnSi. Also the dynamics, i.e., the characteristic relaxation times and their temperature dependence, resembles the behavior of MnSi.

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FIG. 1. Phase diagram of  $Mn_{1-x}Fe_xSi$  deduced from our previous SANS and susceptibility measurements [32,33]. The pink shaded area indicates the precursor phase, P, and PM stands for the paramagnetic phase. The blue line indicates the transition to the ordered helimagnetic phase with long (LR) and short (SR) range periodicity respectively. The temperatures T' have been determined as shown in Figs. 2(b) and 2(c).

In order to understand these results we discuss the evolution of magnetic interactions with doping and also compare  $Mn_{1-x}Fe_xSi$  with MnSi under pressure. Our analysis brings us to the conclusion that in  $Mn_{1-x}Fe_xSi$  doping introduces frustration. We argue that this effect explains both the destabilization of long range helimagnetic periodicity at  $x^*$  as well as the robustness of the precursor phase and its expansion with increasing *x*.

#### **II. EXPERIMENTAL RESULTS**

The samples were grown using the Bridgeman method. For x = 0.09, 0.11 we measured the same single crystals as in our previous SANS study [32]. For x = 0.14 and in order to compensate for the neutron intensity losses at this high Fe doping we chose a large polycrystalline and racemate polycrystalline sample, from the same batch as the samples of our previous studies [32,33]. On one hand, this choice is not problematic, since at this composition the magnetic Bragg peaks have disappeared and the magnetic correlations are no longer pinned to the lattice. On the other hand, as it will be explained below, the polarized neutron scattering from this sample cannot be used to determine the degree of chirality of the magnetic correlations.

The measurements were performed at the IN11 spectrometer, of the Institut Laue Langevin, using the paramagnetic NSE configuration and an incoming wavelength of 0.55 nm. For each sample we determined the polarized neutron scattering and the NSE spectra at the respective maxima of the magnetic scattering intensity. These occur at the scattering vector values  $Q = \tau = 0.58$ , 0.68 and 0.88 nm<sup>-1</sup> leading to helical pitches  $\ell = 10.8$ , 9.2, and 7.1 nm, for x = 0.09, 0.11, and 0.14 respectively, in good agreement with our previous SANS results [32].

By exploiting the polarization analysis capabilities of the experimental setup we obtained an accurate determination of the magnetic scattering and of the chiral fraction of the magnetic correlations [38–43]. The results, shown in



FIG. 2. Temperature dependence of the helimagnetic SANS intensity (a) and the left-handed chiral fraction of the scattering [(b) and (c)]. For the sake of comparison between the results of the three compositions investigated here and the parent compound MnSi (data from [38,39]), the abscissa is the scaled temperature difference  $\beta$  ( $T - T_C$ ), with  $\beta = 2.5$  for MnSi and  $\beta = 1$  otherwise. In this way is is possible to account for the broadening of the precursor phase found for the doped samples.  $T_C = 7.8$ , 5, and 2.5 K for x = 0.09, 0.11, and 0.14 respectively. Panel (c) shows a close-up view of the green rectangle area shown in (b). The dotted lines in (c) illustrate the determination of T'. For T < T' the scattering of MnSi, Mn<sub>0.91</sub>Fe<sub>0.09</sub>Si, and Mn<sub>0.89</sub>Fe<sub>0.11</sub>Si reaches full left-handed chirality within at least two times the error bars.

Figs. 2(b) and 2(c), highlight the abrupt change of behavior at  $x^*$ .

For x = 0.09, i.e., for  $x < x^*$ , a jump of intensity marks the onset of the helimagnetic Bragg peaks and the first-order phase transition at  $T_C = 7.8$  K. This jump disappears for  $x > x^*$ . However, for x = 0.11 the precursor correlations build up with decreasing temperature in the same way as for x = 0.09. Consequently, the precursor phase builds up in a similar way on both sides of  $x^*$ , a result that is hard to reconcile with the For x = 0.14, the evolution of intensity with temperature is much slower and can only be superimposed with data from lower dopings assuming a negative  $T_c$ .

Figures 2(b) and 2(c) show that the scattering from both single crystalline samples, with x = 0.09 and 0.11, is fully chiral at low temperatures. Unpublished results on single crystals indicate that the correlations remain chiral for dopings even higher than  $x_C$  [45]. However, the scattering from our 14% polycrystalline sample was achiral. This brought us to the conclusion that it is a racemate, combining grains with different structural chiralities, because in this system structural and magnetic chiralities are coupled [46]. Thus, the polarized neutron scattering from this sample does not reflect the chirality of magnetic correlations, and for this reason this concentration is not included in Figs. 2(b) and 2(c).

For x = 0.09 and 0.11, full chirality extends up to  $T' \sim T_C + 2.5$  K, thus well above  $T_C$ . The temperature interval  $T' - T_C$  is ~2.5 times broader than in MnSi, where full chirality is found up to  $T' \sim T_C + 1$  K [38,39]. In order to account for this difference when comparing the behavior of the pristine and doped samples, the abscissas in Fig. 2 are the scaled temperature differences  $\beta (T - T_C)$ , with  $\beta = 2.5$  for MnSi and  $\beta = 1$  otherwise. In these plots Mn<sub>0.91</sub>Fe<sub>0.09</sub>Si reproduces the behavior of MnSi, for both the intensity and the chiral fraction.

Further insight in the effect of chemical doping on the magnetic behavior is provided by the intermediate scattering functions  $I(Q = \tau, t)$ , determined by NSE spectroscopy and shown in Fig. 3. The spectra follow an exponential decay that can be fitted by the function  $I(Q = \tau, t) = (1 - a_{el}) \exp(-t/t_0) + a_{el}$ , where  $t_0$  is the characteristic relaxation time and  $a_{el}$  the elastic fraction of the scattering. This behavior contrasts with that of other disordered helimagnets, such as Fe<sub>0.7</sub>Co<sub>0.3</sub>Si [47,48] or Zn doped Cu<sub>2</sub>OSeO<sub>3</sub> [49], where strong deviations from exponentiality have been reported.

The exponential relaxation rules out a spin-glass-like ground state, the footprint of which would have been a stretched exponential decay [50,51]. Furthermore, the NSE spectra become completely elastic at the base temperature, a result which excludes the spin liquid scenario suggested for  $x > x^*$  [35–37]. The elastic fraction, depicted in Fig. 4(a), reflects the evolution of the scattered intensity shown in Fig. 2(b). The change at  $T_C$  is almost step-like, characteristic of the first-order phase transition, for MnSi and Mn<sub>0.91</sub>Fe<sub>0.09</sub>Si. On the other hand, for  $x > x^*$ ,  $a_{el}$  increases gradually with decreasing temperature, as also found in the disordered helimagnet Fe<sub>0.7</sub>Co<sub>0.3</sub>Si [47].

The characterisitic relaxation times,  $t_0$ , depicted in Figs. 4(b) and 4(c), do not change with doping and are comparable to those of MnSi [38,39]. Their values vary between 0.1 and 2 ns, leading to characteristic energies,  $\hbar\omega$ , between 6.58 and 0.33  $\mu$ eV. These values are much lower than those reported by a previous study [30], which, however, suffered from a low Q and energy resolution. The energies found here correspond to temperatures between 80 and 4 mK. Consequently, these are classical fluctuations, with  $\hbar\omega \ll kT$ , not the quantum fluctuations discussed in the literature [35–37]. Thus, classical critical slowing down governs the dynamics





FIG. 3. Intermediate scattering functions, determined by neutron spin echo spectroscopy, of  $Mn_{0.91}Fe_{0.09}Si$  (a),  $Mn_{0.89}Fe_{0.11}Si$  (b), and  $Mn_{0.86}Fe_{0.14}Si$  (c). The lines represent fits to an exponential decay (see text).

of helimagnetic correlations for  $x > x^*$ , masking the quantum criticality associated with the putative QCP at  $x_C$ .

### **III. DISCUSSION**

Our results show that Fe doping affects the first-order transition and the long range helical periodicity in a very different way than the precursor phase. While the former disappear abruptly at  $x^*$ , the latter expands and persists up to higher dopings. Also the characteristic relaxation times of the fluctuations are comparable to those found in MnSi. In order to understand these results and in particular the robustness of the precursor phase, we adopt the Dzyaloshinskii model for cubic noncentrosymmetric ferromagnets [52–54], which leads to the free energy per unit cell [55]

$$\mathcal{E} = \frac{Ja^2}{2} \sum_{i=x,y,z} \partial_i \boldsymbol{m} \cdot \partial_i \boldsymbol{m} + Da \, \boldsymbol{m} \cdot \boldsymbol{\nabla} \times \boldsymbol{m} - a^3 \mu_0 M \boldsymbol{m} \cdot \boldsymbol{H} + \mathcal{E}_a,$$
(1)



FIG. 4. Evolution with temperature and doping of the elastic fractions  $a_{el}$  and the characteristic times  $t_0$  deduced from fitting an exponential decay to the NSE spectra of Fig. 3 (see text). In (a) and (c) the evolution of  $a_{el}$  and  $t_0$  is compared with that of parent compound MnSi (data from [38,39]) and for this reason the abscissa is, as in Fig. 2, the scaled temperature difference  $\beta (T - T_C)$ , with  $\beta = 2.5$  for MnSi and  $\beta = 1$  otherwise.

with J the ferromagnetic Heisenberg exchange, D the Dzyaloshinskii-Moriya (DMI) interaction, m the unit vector in the direction of the magnetization, M the magnetization, a the lattice constant, and  $\mathcal{E}_a$  the magnetic anisotropy energy. The ground state consists of helical spirals with a periodicity of  $\ell = 2\pi a J/D$  propagating along specific crystallographic directions fixed by magnetic anisotropy. A magnetic field strong enough to overcome the anisotropy aligns the spirals towards its direction and tilts the magnetic moments by an angle given by  $\cos \theta = H/H_{C2}$ , where  $H_{C2}$  is the critical field above which the homogeneous or field-polarized state sets in. The model of Eq. (1) leads to  $\mu_0 M_0 H_{C2} = D^2/Ja^3$ , with  $M_0$ the volume magnetization at  $H_{C2}$ . Therefore, at low temperatures, the pitch of the helical spirals can be written as  $\ell =$  $2\pi a \sqrt{J/(a^3 \mu_0 M_0 H_{C2})}$  and, by assuming that J is given by the Curie-Weiss temperature,  $T_{CW}$  (mean field approximation), it can be derived from quantities that have been determined experimentally (see, e.g., Table 1 of [33]). In particular the lattice constant, a, does not change significantly with temperature [11] but varies linearly with x, between 0.4565 nm for MnSi and 0.449 nm for FeSi [46].

By combining this experimental input we calculate the values of  $\ell$  that are depicted in Fig. 5 and vary very little with doping. For the parent compound MnSi the relative difference between the experimental and calculated values of  $\ell$  does not exceed 6%, which gives confidence to our approach. For the sake of comparison Fig. 5 also shows the pressure dependence of  $\ell$  for MnSi (base temperature data from Fig. 2(e) of [7]), which is weak and follows the trend of the calculated values. This contrasts with the behavior of Mn<sub>1-x</sub>Fe<sub>x</sub>Si, where  $\ell$  decreases rapidly with increasing Fe doping. At x = 0.11, the highest Fe concentration where it was still possible to determine both  $M_0$  and  $H_{C2}$ , the experimental value of  $\ell$  is half the one expected from the model of Eq. (1).

An even more drastic reduction of the helical periodicity was recently found in MnSi<sub>1-x</sub>Ge<sub>x</sub> [56,57] and has been attributed to frustration [58]. The effect of frustration can be explained by the interplay between the DMI and higher-order derivatives terms in Eq. (1). These were the original skyrmion stabilization mechanism considered by Skyrme [59] and lead to an attracting skyrmion-skyrmion or soliton-soliton interaction [60]. The numerical simulations in [58] reveal a drastic decrease of  $\ell$  in the presence of an additional antiferromagnetic exchange  $J_{AF}$  term in Eq. (1). This reduction occurs even for weak frustration and, although it was not possible to derive an analytical expression of general validity, the limit  $\ell \gg a$ , leads to [58]

$$\ell \approx 2\pi a \frac{J}{D} \left( 1 - 4 \frac{J_{AF}}{J} \right). \tag{2}$$

Following this approximation for Mn<sub>0.89</sub>Fe<sub>0.11</sub>Si, where  $\ell/a \approx 18$ , the 50% pitch reduction leads to  $J_{AF}/J \approx 1/8$ . This relatively low ratio points to weak frustration and is consistent with the weak disorder seen by electron spin resonance [61] and the exponential relaxation of the NSE spectra.

Antiferromagnetic interactions may result from oscillatory Ruderman-Kittel-Kasuya-Yosida (RKKY) exchange between localized magnetic moments, as suggested by the analysis of the ordinary and anomalous Hall effects [62]. These achiral interactions increase with increasing x and this effect would explain the destabilization of the helical periodicity at  $x^*$ . It was suggested that RKKY interactions arise from the modification of the electron and hole concentrations by the substitution of Mn by Fe [62]. Thus, similarly to MnSi under pressure, the modification of the electronic state drives



FIG. 5. Evolution of the helical pitch as a function of Fe doping, in the case of  $Mn_{1-x}Fe_xSi$ , or as a function of pressure, in the case of MnSi (base temperature data from [32] and [7] respectively). For the sake of comparison the abscissa is either the relative doping  $(x - x^*)/x^*$  for  $Mn_{1-x}Fe_xSi$  or the relative pressure  $(p - p_C)/p_C$  for MnSi. The red dots have been calculated for  $Mn_{1-x}Fe_xSi$  using the model of Eq. (1).

the destabilization of the long range helical order and of the sharp first-order phase transition at  $T_C$ . However, the specific microscopic mechanisms are different for the two systems, as highlighted by the evolution of the helical pitch  $\ell$  shown in Fig. 5. Frustration and disorder are important for  $Mn_{1-x}Fe_xSi$ , in contrast to MnSi under pressure.

Frustration can stabilize spiral and skyrmion periodic states [60,63] but, unlike DMI, it does not impose the chirality, i.e., it does not impose a sense of rotation. This leads to rich phase diagrams, with phases that are impossible in chiral magnets. The competition between frustration and DMI, however, has only been touched upon in the literature [58,64,65] and only in the low-temperature regime with a fixed magnetization modulus. In fact, the influence of frustration on the phase transition and the precursor states in chiral magnets is a most challenging and unresolved problem, which we approach based on general considerations.

Near the ordering temperatures, the functional of Eq. (1) must be supplemented by the Landau expansion in powers of the magnetization [54]:

$$W_0 = A (T - T_0)M^2 + B M^4, (3)$$

where B > 0 and  $T_0$  is the ferromagnetic ordering temperature, i.e., the Curie temperature in the absence of DMI. Under the influence of temperature and of an applied magnetic field, this term enables variations of the magnetization amplitude, which lead to sizable effects in the vicinity of  $T_C$ , the transition temperature in the presence of DMI. In the precursor region of chiral magnets, the chiral twisting is accompanied by strong longitudinal modulations of the magnetization and its interplay with rotational modes [17,66–68]. Thus, anomalous spin textures can be expected, such as a staggered half-skyrmion lattice [17]—a close analog of the square lattice of merons in frustrated magnets [69]. Even in the absence of frustration these precursor modulated states may have both senses of magnetization rotation. However, the modulus increases or decreases depending on whether the rotation adopts the correct or the wrong rotational sense [67,70] with the correct chirality eventually dominating close to  $T_C$ , as shown in Fig. 2. This mechanism induces fan-like oscillations of isolated skyrmions [67] and solitons [70] and leads to their attracting interaction with potentials containing a plethora of local minima at different mutual distances.

An increasing frustration will amplify these precursor modulations and will thus enhance and expand the precursor region, as is indeed the case for  $Mn_{1-x}Fe_xSi$  [28–30,32,33].

# **IV. CONCLUSION**

To conclude, our results shed light on the evolution of helimagnetic correlations and fluctuations in  $Mn_{1-x}Fe_xSi$  for  $x < x_C$ . On one hand, our observations rule out both the spin liquid and the quantum fluctuations hypothesis for  $x > x^*$ , which cannot be associated with a quantum critical point. On the other hand, our analysis indicates that, with increasing doping, frustration increases, which we attribute to increasing competition between chiral DMI and achiral RKKY interactions. We argue that frustration explains the expansion of the precursor phase for  $x < x_C$  and the destabilization of the long range heilmagnetic periodicity at  $x^*$ .

More generally, both frustration and DMI can stabilize spiral and skyrmion periodic states, and the competition between the two can lead to rich phase diagrams with new phases and spin configurations, providing a fertile ground for future developments.

Data from this work are available online [71].

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